

A Quest for Missing Proteins: update 2015 on Chromosome-Centric Human Proteome Project

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** The authors want to pay tribute to Juan Pablo Albar, a friend and recognized scientist, who passed away in July 2014 in Madrid during preparations of the HUPO Congress 2014.

TITLE RUNNING HEAD: A quest for missing proteins

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ABBREVIATIONS

ASV	Alternative Splicing Variant
ATAQS	Automated and Targeted Analysis with Quantitative SRM
B/D HPP	Biology/Disease driven Human Proteome Project
CAPER	Chromosome Assembled human Proteome browser
CHCPC	Chinese Human Chromosome Proteome Consortium
C-HPP	Chromosome Centric Human Proteome Project
CPTAC	Clinical Proteomic Tumor Analysis Consortium
DHS	DNase I hypersensitivity
EBI	European Bioinformatics Institute
ENCODE	Encyclopedia of the DNA Elements
ETD	Electron Transfer Dissociation
ETHcD	combination of electron-transfer dissociation (ETD) and higher-energy collision dissociation (HCD)
FDR	False Discovery Rate
GEO	Gene Expression Omnibus
GPCR	G-protein coupled receptors
gpmDB	Global Proteome Machine database
GSC	<i>Glioma</i> stem cells
HPA	Human Protein Atlas
HPPP	Human Plasma Proteome Project
HTML	HyperText Markup Language
HUPO	HUMAN Proteome Organisation
ISBER	International Society for Biological and Environmental Repositories

JSON	JavaScript Object Notation
MATF	Monash Antibody Technologies Facility (Monash University)
MS	Mass Spectrometry
NIST	National Institute of Standards and Technology
ORF	Open Reading Frame
PASSEL	PeptideAtlas SRM Experiment Library
PE	Protein Existence
PrESTs	Protein Epitope Signature Tags
PSM	Peptide Spectrum Match
REST	Representational State Transfer
SAV	Single Amino acid Variant
SILAC	Stable Isotope Labeling by Amino Acids
SISCAPA	Stable Isotope Standard Capture with Anti-Peptide Antibodies
SOP	Standard Operating Procedure
SPARQL	Protocol and RDF Query Language
SPE	Solid Phase Extraction
SRM	Single Reaction Monitoring
SWATH-MS	Sequential Window Acquisition of all Theoretical Fragment ion Mass Spectrometry
TPP	Trans Proteomic Pipeline
XML	Extensible Markup Language

WEB LINKS

BioPortal	http://bioportal.bioontology.org/
C-HPP Web	http://www.c-hpp.org
C-HPP Wiki	http://c-hpp.webhosting.rug.nl/
dasHPPboard	http://sphppdashboard.cnb.csic.es/
Disease Ontology	http://disease-ontology.org/
neXtProt REST service	www.nextprot.org/rest/
Protannotator	http://biolinfo.org/protannotator
Ontology Lookup Service	http://www.ebi.ac.uk/ontology-lookup/

Abstract

This paper summarizes the recent activities of the Chromosome-Centric Human Proteome Project (C-HPP) consortium, which develops new technologies to identify yet-to-be annotated proteins (termed “missing proteins”) in biological samples that lack sufficient experimental evidence at the protein level for confident protein identification. The C-HPP also aims to identify new protein forms that may be caused by genetic variability, post-translational modifications, and alternative splicing. Proteogenomic data integration forms the basis of the C-HPP’s activities; therefore, we have summarized some of key approaches and their roles in the project. We present new analytical technologies that improve the chemical space and lower detection limits coupled with bioinformatics tools and some publicly available resources that can be used to improve data analysis or support the development of analytical assays. Most of this paper’s contents have been compiled from posters, slides, and discussions presented in the series of C-HPP workshops held during 2014. All data (posters, presentations) used are available at the C-HPP Wiki (<http://c-hpp.webhosting.rug.nl/>) and in the supporting information.

Introduction

Proteins such as those acting as enzymes, regulatory proteins, transporters, and receptors are the active macromolecules of human biology and are thus central to understanding biological molecular processes. Understanding the diversity and complexity of these biological molecular interactions is a central focus of bio-medical research today. It is important that all protein forms of human genes are eventually studied so that their biological functions and roles in healthy and disease states can be determined [**posters 17 and 18 in Table 1**]¹⁻³. Proteins cannot be amplified and are chemically much more heterogeneous than DNA and RNA. Their analysis therefore represents a much more significant analytical challenge.

To meet this challenge, the Human Proteome Organization (HUPO) announced in 2010 at the HUPO Congress in Sydney, Australia, the formation of the Human Proteome Project (HPP) to sequentially catalogue the protein products of human genes, both to identify proteins that have little or no evidence at the protein level, termed “missing proteins”^{4, 5}, and to discover and characterize protein sequence variability with genetic origin and post-translational modifications (PTM) of known proteins. The Chromosome-Centric Human Proteome Project (C-HPP)⁴⁻⁶ is a large multidisciplinary international effort to identify all human protein forms and catalogue them on the basis of the chromosome location of their coding genes. In the C-HPP, one national or multinational team is responsible for the identification and annotation of protein products of the genes in each chromosome. Evidence at the protein level means that a protein has been detected, preferably by mass spectrometry (MS) and preferably from multiple peptides unique to the protein observed in multiple experiments and possibly in diverse biological sample types. Multiple data resources help to provide information on protein evidence.

The sharing of data, protocols, and other electronic resources such as proteome annotations is crucial in the C-HPP and proteomics community to enable the reuse of collected information (e.g., in form of high-quality spectral libraries for spectral library identification or the development of selected reaction monitoring [SRM] assays) and to develop and improve new data analysis protocols. The current status and developments in the shared proteomics data and proteome knowledge by the main stakeholders are presented below. An overview of current bioinformatics resources and the data flow to support the creation of complete part lists of the human proteome is shown in **Figure 1**.

The ProteomeXchange⁷ consortium, led by PRIDE^{8, 9} at the European Bioinformatics Institute (Hinxton/Cambridge, UK) and by PeptideAtlas¹⁰⁻¹² at the Institute for Systems Biology (Seattle, Washington, USA), is devoted to the standardization of data submission and dissemination of MS-based proteomics data and to the promotion of public sharing of proteomics data in the public domain.

In addition, it promotes the use of community data standards developed by the Proteomics Standards Initiative (PSI). As such, ProteomeXchange resources store original MS datasets, containing at least the raw MS data accompanied by the processed results (peptide and protein identifications, but possibly quantitative information as well) and by suitable experimental and technical metadata. By November 2014, ProteomeXchange resources stored approximately 1500 datasets (~50% of which are publicly available) from a wide variety of sources, with humans being the most-represented species. Once the datasets are made publicly available, they are usually reprocessed by PeptideAtlas¹⁰⁻¹² using the Trans Proteomic Pipeline (TPP)¹³⁻¹⁷ and by the Global Proteome Machine Database (gpmDB)^{18, 19} using the X!Tandem^{20, 21} database search tool. Basically, everyone in the community can reanalyze the raw data available in ProteomeXchange for different purposes.

To provide a comprehensive view of the human proteome and its diversity, neXtProt^{22, 23} is adding information at the genomic, transcriptomic and proteomic levels to the corpus of information available in UniProtKB²⁴⁻²⁶. In particular, neXtProt integrates genomic variation data from dbSNP²⁷ and COSMIC²⁸, transcriptomic data from BGee²⁹, antibody-based protein evidence from the Human Protein Atlas (HPA)^{30, 31}, MS-based information from PeptideAtlas¹⁰⁻¹², three-dimensional structural information from the Protein Data Bank³² and various PTM information from manually curated literature. Based on this combined information, neXtProt attributes a protein existence (PE) level to each entry originally defined in UniProtKB. The PE1 level (experimental evidence at the protein level) denotes entries with credible evidence by protein expression and identification by MS, immunohistochemical analysis, three-dimensional structure, or amino acid sequencing. The PE2 level (experimental evidence at transcript level) refers to proteins with transcript expression evidence but without evidence of protein detection. The PE3 level (protein inferred from homology) is attributed to proteins without human protein or transcript evidence but with strong evidence on homologous protein in another species. The PE4 level (protein predicted) is for proteins that are hypothesized from gene

models, and the PE5 level (protein and gene uncertain) refers to “dubious” or “uncertain” genes that at one time seemed to have some protein-level evidence but have since been deemed doubtful. The PE5 category generally corresponds to pseudogenes or non-coding RNAs according to the protein annotation from different resources (HGNC³³, RefSeq^{34, 35}, HAVANA, CCDS³⁶⁻³⁸, UniProtKB/Swiss-Prot³⁹). Among the 643 entries in the PE5 category in neXtProt in August 2011, 119 have already become obsolete in UniProtKB and have been deleted from neXtProt, 13 have been upgraded to the PE1 category due to manual curation of publications and/or convincing proteomics data, and 11 have been upgraded to the PE2 or PE3 categories. Based on these numbers, one can estimate that less than 5% (< 30) of the remaining PE5 proteins in neXtProt are true proteins. Given this low probability, any MS identification of PE5 proteins must be carefully checked. Proteins in the PE2-4 categories are awaiting experimental confirmation at the protein level and are called “missing proteins” in the context of C-HPP.

One of the primary tasks of the C-HPP is to determine why no protein products have been identified for certain genes showing open reading frame for translation, i.e., genes coding for the so-called “missing proteins with no or poor protein evidence.” There are five main reasons for the existence of “missing proteins” (**Figure 2**)⁴⁰. (1) The current mainstream proteomics technology cannot identify them, possibly because of the low abundance of the proteins, because the sequences do not contain tryptic cleavage sites or generate peptides which can uniquely identify the proteins, or because the protein digestion results of peptides that are lost during the sample preparation and analysis. (2) They are expressed only in rarely studied tissues or cell types, or are expressed only as a result of a stimulus or perturbation. (3) They are not expressed at all and are part of the silent information of the human genome. (4) They reflect erroneous annotation of the genome, which results in incorrectly predicted protein sequences. (5) Many highly homologous proteins or proteins with large sequence variability are missed or not counted due to the parsimonious protein assembly of shotgun MS/MS protein

identification or due to large sequence variability such as immunoglobulins. PeptideAtlas and gpmDB select only one “representative protein” among highly homologous members of protein families⁴⁰ when the available sequence coverage cannot distinguish these related proteins (see the Cedar scheme in Farrah et al.⁴¹ and **Figure S1** in the supporting information).

When the C-HPP began (2012), it was announced that no satisfactory evidence existed at the protein level for 6568 (33%) of the 20,059 protein-coding genes⁴². NeXtProt released a new version as of September 19, 2014, that contains 20,055 entries, among which 16,491 are PE1 proteins, 2948 lack protein evidence (PE2, PE3, PE4), and 616 are dubious (PE5). Hence, there is still insufficient evidence at the protein level for approximately 15% (if we exclude PE5) or 18% of the human proteome. It is encouraging to see that the number of missing proteins has been reduced considerably since the initial assessment by the C-HPP in 2012.

This paper presents an overview of the new technologies and new resources that have been used during the last 4 years by members of the C-HPP consortium and others to identify missing proteins. Most of the contents described here have been obtained from data presented during several HUPO workshops in 2014, including the 9th, 10th and 11th C-HPP Workshops in Busan, South-Korea (26 March), Bangkok, Thailand (9 August) and Segovia, Spain (9 October), respectively, and during the C-HPP and HPP sessions of the HUPO 2014 Congress in Madrid, Spain (5-8 October).

Proteogenomics

Analytical technologies and bioinformatics are the key components for the identification and quantification of proteins in a complex biological sample. The current workhorse of proteomics analysis is shotgun LC-MS/MS, typically using a C18 stationary phase and acetonitrile/water eluent pairs, resulting in sequence coverage typically lower than 30% for identified proteins. Additionally, most of the collected MS/MS spectra contain gaps in the fragment ion series, thus preventing de novo

peptide sequence spectra interpretation and identification⁴³. The most widely used approach for protein identification is database searching, which requires a list of protein sequences that are expected to be present in the analyzed samples. UniProtKB is the most frequently used protein sequence database. It has two main components: UniProtKB/SwissProt (which contains manually curated sequences) and UniProtKB/TrEMBL⁴⁴ (which contains computationally generated records from DNA sequences that have not been manually curated). Canonical sequences are used to represent the most prevalent sequences that are most similar to those of other species and in which the length or amino acid composition allows the clearest description of protein domains, splice isoforms, polymorphisms, and PTMs. UniProtKB contains some degree of protein isoform information and sequence variation due to genetic variability, but it is far from complete. This results in a low level of identification of peptides that arise from alternative splicing, coding non-synonymous single-nucleotide polymorphisms (SNPs), and single amino acid variants (SAVs) due to RNA editing^{45, 46}. Therefore a proteogenomic approach using DNA and mRNA data to build a protein database that contains all genetic variability or sample-specific protein sequence information is becoming more and more popular and has contributed to the identification of new protein forms⁴⁷. Conversely, peptide-level data can serve to fill gaps or correct errors in the DNA and RNA databases^{47, 48}. There are two main proteogenomic approaches: (1) a protein sequence database is constructed from publicly available databases that contain sequence information with genetic variability such as dbSNP^{27, 49} or H-INVDB⁵⁰, or (2) a customized protein sequence database is constructed from annotated DNA and mRNA transcript data obtained from the same sample⁵¹⁻⁵³. However, proteogenomic analysis generally results in a larger database than does using the UniProtKB canonical sequences, and should be followed through false discovery rate (FDR) analysis at both the peptide and protein levels^{13, 47}, especially when the database search is performed in multiple steps^{47, 54}. PeptideProphet⁵⁵ from TPP and Percolator⁵⁶⁻⁵⁸ can be used for FDR calculations for peptide spectrum matches (PSMs). ProteinProphet⁵⁹ and MAYU¹⁷ (both from TPP) serve to estimate

FDR for protein inference. PeptideShaker^{60, 61} provides a solution for both PSM and protein FDR calculation. The statistical power of PSM in separating correct and incorrect PSM distributions can be enhanced by including the measurable and predictable physico-chemical properties of peptides in addition to m/z , such as the retention time in liquid chromatography⁶² or the high-resolution isoelectric point⁶³. Identification of missing proteins can be enhanced by identifying cell lines and tissue samples with transcriptomics evidence [**poster 12**], analyzing samples of different ages, and including samples acquired under special stress conditions and biological perturbations.

A general drawback of bottom-up shotgun LC-MS/MS approaches is that complete protein forms cannot be reconstituted from peptide fragments. A top-down approach that allows the peptide-protein interference problem to be avoided may provide a solution for determination of the accurate distribution of whole protein forms, also called proteoforms⁶⁴. Proteoforms are the most recent nomenclature of protein forms introduced by the Top Down Proteomics Consortium, which “designates all of the different molecular forms in which the protein product of a single gene can be found, including changes due to genetic variations, alternatively spliced RNA transcripts and post-translational modifications”. The relationship of the proteoform terminology to the UniProt canonical sequences and other protein sequence variability or modifications is shown in **Figure S2** in the supporting information. Importantly, unlike bottom-up protocols in which detailed information on PTMs and sequence variants is compromised because of enzymatic digestion, intact proteins are analyzed in top-down approaches, which allows the unequivocal identification and location of specific modifications. However, they require relatively pure protein samples, they are restricted to proteins of less than 30 kDa, the available fragmentation spectra are often far from complete, and the obtained complex spectra are often difficult to interpret⁶⁴⁻⁶⁷.

Analysis of mRNA has an advantage in that sequences can be amplified to provide nearly complete sequence coverage using current RNA sequencing technologies. The challenge is to accurately annotate

the resulting raw DNA and RNA data, which is generally performed using the Ensembl genome browser⁶⁸. Ensembl contains a reference genome and includes annotation from the Encyclopedia of the DNA Elements (ENCODE)^{69, 70} which is a “comprehensive parts list of functional elements in the human genome, including elements that act at the protein and RNA levels, and regulatory elements that control cells and circumstances in which a gene is active.” However, protein-coding gene annotations such as GENCODE⁷¹ are based on the protein sequences stored in public databases such as UniProtKB or NCBI RefSeq³⁵ and gene models that predict the long open reading frames (ORFs) that are most likely to code a protein, which can lead to errors in the annotation of these databases. Therefore, besides revealing protein forms due to genetic variability, a proteogenomic approach can help to confirm the existence of the 616 dubious human proteins currently annotated as PE5 in neXtProt. It can also support identification of new ORFs and translated non-coding mRNA, or redefine the starting and ending parts of protein coding regions, as reported by Kim et al^{1, 72}. However when protein identification is performed exclusively with a translated mRNA sequence the much shorter half-life of mRNA compared to proteins^{73, 74} should be taken into account in the integration of proteogenomics data. The half-life difference between these two molecular species could result in proteins without mRNA when proteins and mRNA are measured in the same sample and at a single time point. Time series sampling could be used to overcome this issue, when it is possible. This is the case for cell cultures, blood or animal experiments, or tissues for which multiple samples are available from the same specimen at different time. For other cases, the use of a combined databases from translated mRNA sequences and the UniProt database is an option for the detection of proteins with a half-life much longer than that of mRNA.

Translating mRNA, which is directly upstream of protein expression, thus serves as a useful resource for protein identification⁵³. Wang et al.⁷⁵ performed the first translated mRNA sequencing (RNC-seq) in human lung cancer cell lines and observed an improved correlation of RNC-mRNA abundance with

translated protein when the RNC-mRNA length was taken into consideration. The same group showed that the genes with translation evidence represent an improved reference for the identification of proteins, the detection of sequence variations (SAV, RNA editing and alternative splicing), and integration of the MS data⁵³.

Furthermore, missing proteins with mRNA evidence and more stringent conditions with ribosome-bound mRNA (RNC-mRNA) evidence are most probably translated, but the current proteomics technology does not allow their detection because of a restricted chemical space or because the detection sensitivity is not sufficient. According to the presentation from Zhang et al. (submitted manuscript) at the C-HPP workshop during the HUPO 2014 Congress in Madrid, ~5% of transcribed mRNAs are typically not translated in a single cell line, and these non-translated mRNAs are highly cell-type specific and/or tissue specific. This allows the focus to be placed on missing proteins in samples with translation evidence and the development of targeted SRM assays and specific sample preparation methods, e.g., the use of antibody enrichment of missing proteins for low abundant peptides or the use of different proteases when missing proteins do not contain identifiable unique tryptic peptides with the detected tryptic peptide set.

An example of a proteogenomic study in which translated mRNA analysis, proteomics data integration, and the use of antibodies were performed to enrich low abundant proteins was presented by Chang et al.⁵² from the Chinese Human Chromosome Proteome Consortium covering chromosomes 1, 8, and 20. In their study, three hepatocellular carcinoma cell lines (Hep3B, HCCLM3, and MHCC97H) were submitted for mRNA and RNC-mRNA analysis and to comprehensive analysis with deep proteomics and antibody-enriched transcription factor proteomics. Based on the integrated data, they concluded that only 50.2% of the protein-coding genes with translation evidence were found in the proteomic data. This result is comparable to that of a previous study on the RNC-mRNA and MS data of Caco-2 cells: 52.6% of the proteins with translation evidence were missing from the LC-MS/MS data acquired

from institutions⁵³. The inability to detect certain proteins by LC-MS/MS was most probably a result of the translation control mechanisms and analytical limitations of MS-based shotgun identification of peptides and proteins. This warrants a survey of missing proteins in other resources and strategies, such as detergent-insoluble fractions of cell/tissue lysates and forced gene expression through epigenetic manipulations.

Integrating alternatively spliced transcripts with proteomics information allows the study of the transcriptional regulation of proteins in both healthy and diseased tissue [**poster 8** and **13**]. This effect was shown by Menon et al.⁷⁶, who integrated RNA-seq and proteomics data as part of the chromosome 17 team and identified more than one splice variant for each of 1167 genes expressed in at least one of three breast cancer cell line models ERBB2+SKBR3, ERBB2+ SUM190, and EGFR(ERBB1)+SUM149, of hormone receptor–negative breast cancers. Their data analysis showed high differences between alternative splicing distributions in the three different cell lines, which were distinctively enriched for different key cell functions such as amino acid and sugar metabolism, caspase activity, and endocytosis in SKBR3; aspects of metabolism, especially of lipids in SUM190; and cell adhesion, integrin and ERK1/ERK2 signaling, and translational control in SUM149. In **poster 20**, Menon and Omenn presented findings of recurrent non–canonical splice variants of interesting proteins in 126 triple-negative breast cancer specimens using data available from EBI/PRIDE.

Another dimension of the proteogenomic splice isoform studies was presented in **poster 21** by Li et al, who undertook genome-wide isoform-level protein connectivity analysis. The isoform with the highest connectivity seems to be more highly associated with function than the choice of a canonical protein isoform based on the sequence length or the abundance of the isoform, which are the methods commonly used in established databases. The genome-wide isoform analysis in mice has been reported⁷⁷ and is under development for humans by the Chromosome 17 team (Li et al., unpublished).

Glioma stem cells (GSCs) isolated from patient tumors possess both stem-like and oncogenic patterns of protein expression and are thus a potential source of missing proteins. The Chromosome 19 team has characterized their expression profiles at both the transcript and protein levels. They analyzed 1382 chromosome 19 genes in GSCs using a transcription microarray and showed that 70-75% of them were expressed in each of the studied cell lines⁷⁸. The customized analyses identified differential gene expression patterns specific to chromosome 19 between subtypes of GSCs. It was found that roughly 20% of the transcripts were differentially expressed in the proneural and classical subtypes in comparison to transcription patterns in human neuronal stem cells⁷⁹. The chromosome 19 transcripts that potentially encoded candidate unidentified ORFs proteins were also investigated; 43 ORFs were represented on the arrays, of which 31 (72%) were expressed in the GSC lines. GSCs are also a source of protein variants. Recently, proteomic searches of high-resolution LC-MS/MS data of GSC protein digests identified 19 SAVs in 17 chromosome 19 proteins^{1,2}. Several of the protein variants may have oncogenic potential and are the subjects of further investigation. Furthermore, the integration of RNA-seq and proteomic data made possible the study of the somatic-proteomic landscape of GSCs, thereby allowing the contribution of new knowledge regarding novel fusion proteins in GSC pathobiology. To summarize the current status of chromosome 19, **Figure 3** shows the numbers of genes, mRNA, and proteins, including the number of “missing” proteins and the number of predicted molecular forms (such as mutant proteoforms) and known PTMs. This study and the preceding studies from the Chromosome 17 team illustrate the potential of involving new protein forms that arise from genetic variability and alternative splicing into the investigation of new biology.

Enlarging the analyzed chemical space

Proteins are composed of 20 amino acids and are known to be modified by more than 300 types of PTMs⁸⁰⁻⁸², embracing a wide chemical space that should be covered by the proteomics analytical approach. In addition, artificial modifications introduced by the sampling protocol need to be

considered. This large chemical space is well covered, but not completely ~~covered~~, by the widely used acetonitrile/water C18 LC-MS/MS protocols. For example, studies in multiple tissues and cells lines performed by the Chinese Human Chromosome Proteome Consortium⁵² showed that hydrophobicity (28%) and a low molecular mass (<30 kDa; 75%) are important physicochemical properties that predict unsuccessful detection of a protein. In contrast, the isoelectric point and half-life do not seem to play important roles in detectability. Unidentified proteins in hepatocellular carcinoma cell lines were enriched in specific cellular processes such as olfaction with non-liver function or mainly localize in the cell membrane, supporting the hydrophobicity-negative bias of the currently dominant method of proteomics analysis. Tissue transcript analysis showed that transcripts for the missing proteins are abundant in the testis. Interestingly, a recent analysis of data in the HPA has shown that more tissue-specific proteins are made in the testis than in any other tissue in the body⁸³. Analysis of the DNase I hypersensitivity of mRNA and RNC-mRNA data suggests that the missing proteins without a detectable signal are relatively enriched in the chromatin regions with low DNase I hypersensitivity (~40% of the missing proteins), which suggests that the specific structure of chromatin can repress the transcriptional process. Chromosome 11 (and to a lesser extent chromosome 19) showed a greater number of missing proteins without transcript evidence, and those missing proteins were densely clustered in several well-defined chromosome regions. One major group of these missing proteins is presumed to have olfaction function.

Missing protein identification can be enhanced by developing specific enrichment methods such as the use of Proteominer beads^{84, 85} and enrichment of protein aggregates (Yang Chen, Yaxing Li, Jiayong Zhong, et al.; manuscript under review in JPR); specific analytical methods for hydrophobic proteins; a specific fractionation method such as the analysis of subcellular fractions [**poster 5, 9 and 19**]; and methods to increase protein sequence coverage (e.g., by using multiple proteases for protein cleavage⁴⁹ or by using a more efficient method of peptide fragmentation such as EThcD⁸⁶⁻⁸⁸). The membrane

subproteome was suggested to be a rich source of missing proteins. A deep sequencing strategy using complementary two-dimensional chromatography with a combination of high-pH reversed phase (RP), strong anion exchange and low-pH RP stationary phases was used to increase the measured dynamic concentration range. The preliminary results of the enriched membrane proteome from the group led by Yu-Ju Chen [**poster 1**] showed that high-pH RP columns enhanced the retention of hydrophobic peptides and increased the identification coverage of the missing membrane proteins (unpublished results).

Lowering the detection limit with targeted SRM, SWATH analysis, ProteomeAnalyzer, and antibody enrichment

SRM assays have been used for decades to quantify small compounds by MS. The laboratory of Ruedi Aebersold has further developed this approach into a standard method for proteomics to enable simultaneous multiplexed quantification of several hundred proteins in complex biological samples with a wide concentration dynamic range. Picotti et al.^{89, 90} showed the power of this method by detecting almost the complete proteome of yeast, covering 4.5 orders of magnitude of the dynamic concentration range. Large-scale application of SRM assays for targeted quantification of long human protein lists required not only the increased speed of the triple quadrupole instruments, but also the creation of such important informatics resources as high-quality spectral libraries (e.g., NIST spectral libraries, SRMAtlas⁹¹), repository of SRM assay results (PASSEL⁹²), a database of ranked peptides and SRM transitions for all proteins in selected proteomes (SRMAtlas⁹¹), and a database of peptides and transitions with quantification calibration curves (SRMQuantAtlas). SRM assay development requires the identification of proteotypic peptides that not only map uniquely to a single protein or isoform but also are readily ionized and can be detected by MS with a high probability. The proteotypic sequence and SRM transitions must be unique to unequivocally identify the protein form among all other protein forms in the human proteome. This task, coupled with the processing and analysis of the acquired data,

is supported by step-specific algorithms and comprehensive bioinformatics pipelines⁹³ to plan SRM assays for missing proteins, such as ATAQS⁹⁴, mQuest⁹⁵, MaRiMba⁹⁶, SMRBuilder⁹⁷, and Skyline⁹⁸. The PeptidePicker tool developed by Mohammed Y et al.⁹⁹ can help to select the most appropriate surrogate peptides for a given protein list in human and mouse proteomes to be used in targeted SRM assays based on the current knowledge of the community as presented in UniProtKB, PeptideAtlas, gpmDB, PRIDE and dbSNP. The tool identified has already reported peptides in online databases for missing proteins, although the quality of the data in these databases varies considerably.

The data-independent sequential window acquisition workflow (SWATH-MS) allows collection of non-targeted fragment spectra by fragmenting large windows of precursor ions (typically 20 to 25 Dam/z). The resulting MS/MS data can be reconstituted from the co-eluted fragment ions with liquid chromatography retention time using deconvolution methods. The SWATH approach also can be seen as a generalization of the SRM approach, in which each detectable fragment ion is measured and can be reconstituted from the acquired data without being restricted to a targeted list of transitions as in SRM. Recently, SWATHAtlas was introduced, which stores a human library of MS/MS spectra acquired on a TripleTOF instrument for 10,000 human proteins¹⁰⁰. This library was obtained from 331 measurements on cell lines, blood, and other human tissues and is intended to be used by PeakView, the OpenSWATH tool¹⁰¹, and other analogous processing software, providing 51% of coverage of canonical UniProtKB/Swiss-Prot³⁹ entries.

Another important resource for the identification of missing proteins is SRMAtlas⁹¹, which contains a high-confidence “gold standard” quality SRM assay for at least one unique peptide for 99.9% of the canonical UniProtKB/Swiss-Prot³⁹ entries. This high coverage was achieved by including MS/MS spectra obtained from a large campaign of production and analysis of synthetic peptides for the complete human proteome. Another source of MS/MS spectra and spectral libraries for phosphorylated and unmodified synthetic peptides is available for assay development¹⁰².

The development and application of SRM assays to complex biological samples is a well-established technology for protein quantification that requires expensive instrumentation and experienced personnel, which limits its utility in replacement of the commonly used western blot analysis for quantification of proteins. Following a planning period led by the HUPO Industrial Advisory Board and a survey of 266 participants, mostly from biology and clinically oriented laboratories, HUPO launched the ProteomeAnalyzer initiative in collaboration with instrument vendors with the goal of developing affordable SRM instrumentation capable of quantifying of 100 to 2000 proteins.

Antibodies are effective reagents for the specific detection and enrichment of missing proteins¹⁰³. The availability of highly specific and validated antibodies is crucial for the detection of low abundant missing proteins and the spatial characterization of their expression pattern in cells and tissues. The implementation of high-throughput production of validated high-affinity monoclonal antibodies using automated production systems will provide renewable resources^{104, 105}. SISCAPA can enhance the sensitivity of SRM analyses by enriching specific peptides¹⁰⁶⁻¹⁰⁸.

The HPA^{31, 109} project has systematically generated affinity purified polyclonal antibodies using proteospecific recombinant protein fragment and Protein Epitope Signature Tags (PrESTs)¹¹⁰. After a rigorous validation scheme, the approved antibodies are used to assess the spatial distributions of the proteins in a multitude of human cells and tissues by immunohistochemical analysis. The November 2014 HPA release (version 13.0) contains more than 13 million images of protein expression patterns generated by the use of 23,968 validated antibodies targeting 16,943 genes. In addition to protein evidence, expression levels, and subcellular localization, the HPA contains mRNA expression levels for the majority of tissues and cell lines involved in the HPA¹¹¹. The resources from HPA are highly valuable for the identification of cell lines and tissues that express missing proteins or for cross-validation of MS or HPA antibody protein evidence. Methods for the use of PrEST antigens as spike-in reagents for quantitative MS were recently demonstrated¹¹². Immuno-SILAC has proved capable of

absolute quantification of proteins in complex samples based on HPA antibodies and stable isotope-labeled PrESTs to allow affinity enrichment before MS analysis and accurate quantification¹¹³.

In a recent collaboration between the HPA group in Stockholm and the high throughput monoclonal antibody facility at Monash University in Melbourne a number of monoclonal antibodies against missing proteins, important signaling molecules and proteins of interest to the Chromosome 7 and 17 groups were generated using the same PrESTs as immunogens, which will allow a direct comparison between monoclonal and polyclonal antibodies raised against the same prEST and generate new reagents for the proteomics community. Interestingly, in some cases it was possible to raise monoclonal antibodies to targets that had failed to generate polyclonals. This finding provides an additional route for completion of the task of generating renewable antibodies to all human proteins using the existing antigen resources. Lambert et al.¹¹⁴ recently showed that coupling affinity enrichment with quantitative MS techniques such as SWATH analysis provides the most sensitive detection method for low abundant missing proteins.

Human sample resources

Human samples are collected and stored in various locations worldwide and are crucial to the C-HPP project and to proteomics and disease research in general. Even if sensitive analytical methods are available to uniquely identify and detect missing proteins, high-quality human samples collected under strict standard operating procedures for collection, processing, and storage must be available to characterize protein expression. Although many countries have recognized this need and have established biobanks for the collection and storage of human samples available from local or regional resources, they have not always been collected under the optimal conditions required for the maintenance of the initial integrity of the protein constituent of samples for proteomics studies. Here, many factors leading to protein degradation need to be identified and addressed by the community. Therefore, as demonstrated by several groups¹¹⁵⁻¹²¹, sample collection and storage protocols should be

assessed and optimized in this respect for each sample type. For the C-HPP initiative, in addition to ensuring the sample quality, it is also important to exchange samples between laboratories in different countries, for which legal and ethical regulations should be in place. To facilitate the exchange of samples, HUPO will join forces with ISBER^{122, 123}, an international organization that has worked out regulatory and ethical protocols and Best Practice guidelines¹²⁴ for such purposes.

Controlled vocabularies and ontologies pioneered by SNOMED¹²⁵ providing standardised anatomical descriptors related to tissue types (BRENDA)^{126, 127}, cell types (Cell Ontology)¹²⁸, and human diseases (DOID, <http://disease-ontology.org/>)¹²⁹ and common descriptions of clinical details, sampling, sample handling, and sample storage data are crucial to effectively compare and search metadata of the samples stored in biobanks and to enable studies that make use of samples from multiple biobanks. Biology and clinically related ontologies are accessible through the Ontology Lookup Service¹³⁰ hosted at EBI (<http://www.ebi.ac.uk/ontology-lookup/>) or at BioPortal (<http://bioportal.bioontology.org/>).

Integration of the HUPO Biology/Disease Human Proteome (B/D HPP) and C-HPP initiatives will be beneficial for both consortia because C-HPP can provide new assays for missing proteins or protein isoforms whose role and function can be immediately studied by B/D HPP teams in the context of health and disease. G-protein-coupled receptors, and especially olfactory receptors, are overrepresented among the missing proteins. This protein family is low abundant and shows highly specific tissue expression, and expression of the approximately 900 human olfactory receptors that are responsible for the detection of odorant compounds is only expected in nasal tissue. An assessment of the number of identified olfactory receptors in Kim et al.⁷² and Wilhelm et al.¹³¹ by Ezkurdia et al.¹³² showed that these two large-scale studies with poor MS/MS spectra identified more than 100 olfactory receptors, despite the fact that they did not include data from nasal tissue. This quality assessment shows the importance of critical error analysis of peptide and protein identification in large-scale data analysis projects. The use of a 1% threshold for FDR limited only to PSM or peptide levels is not

sufficient to provide a high-quality list of identified proteins in large aggregated datasets. Therefore the statistical criteria must be a 1% FDR or better calculated at the protein level for the combined dataset as adopted by PeptideAtlas^{11, 17}. Using a 1% FDR threshold at the PSM or peptide level would result in a large number of misidentified or indistinguishable proteins when analyzing a large amounts of data. These incorrect PSMs map to proteins randomly, which results in a greater FDR at the protein level. Setting an FDR should take into account the number of identified peptides and proteins in large datasets. For example, if a million PSM pass a threshold of 1% FDR, this implies that there are 10,000 false PSMs, and these tend to map to proteins with one peptide per protein, which results in large FDR at the protein level. For datasets from which 3000 proteins are identified, a 1% protein-level FDR implies only 30 incorrect protein identifications. However, for very large datasets from which 15,000 proteins are identified, a 1% protein-level FDR would result in 150 misidentified proteins, which is a considerable number. In this case, lowering the FDR to 0.1% for example, would keep the number of misidentified proteins at more acceptable number of approximately 15. C-HPP will stringently identify olfactory receptors in nasal tissue accompanied with thorough FDR analysis at the PSM, peptide and protein levels.

Bioinformatics resources

High-level bioinformatics support is crucial for the success of the C-HPP initiative and goes beyond the already-listed sequence knowledge bases, MS databases and SRM assay development support, and evaluation pipelines. Many groups have developed Human Proteome Browsers to support the chromosome-centric integration, processing, and visualization of proteogenomic data or MS/MS repositories such as the Gene-Centric Knowledgebase^{133, 134}, GenomeWideDB¹³⁵ [**poster 4**], Human Proteome Map⁷², proteomicsDB¹³¹, gpmDB¹⁸, PeptideAtlas^{11, 12, 91}, HPA^{6, 30, 31, 109}, The Proteome Browser¹³⁶, CAPER^{137, 138} [**poster 2**], and Human Proteinpedia¹³⁹⁻¹⁴¹. These resources are currently

being developed in isolation, which makes it difficult to further interrogate the diverse types of information stored in these resources. With the participation of the major database developers listed previously (**Figure 1**), an initiative to create a Unified Human Proteome Browser [**poster 16**] as an advanced knowledge-mining system was established at HUPO 2014 in Madrid. This builds on the strengths of existing browsers and their development teams to provide a unified platform for further detailed analysis of the acquired proteogenomic data from the perspectives of chromosomes, biology, and disease. This will lead to a better overview of the existing proteogenomic information that can be developed to suit the needs of the global proteomics community and to improve the current standards of data processing, visualization, and interpretation. It will be essential to subject the component resources and their overall performance to comparisons of assumptions, methods, or findings.

The importance of the quality of bioinformatics workflows and use of false-discovery thresholds was demonstrated by Eric Deutsch, who showed that the addition of four large datasets (the CPTAC repository¹⁴² and those of Kim et al.⁷², Wilhelm et al.¹³¹, and Guo et al.¹⁴³) to PeptideAtlas^{11, 12, 91} only increased the amount of level 1 protein evidence for approximately 1365 neXtProt entries using stringent error thresholds of 0.000091 FDR for PSMs, 0.00028 FDR at peptide level, and 0.011 FDR at protein level identification. The successive increments in HumanAll build database of PeptideAtlas from these new large studies were 541, 591, 231, and 2 proteins. gpmDB, PeptideAtlas, and neXtProt each estimated the high-quality protein identifications from Kim et al.³⁹ and Wilhelm et al.⁶⁰ to be about 13,000, not 17,294 or 18,059, as reported. Further scientific scrutiny of the many reasons for these large discrepancies will be desirable, involving all parties, as launched in Madrid.

The proteomics community has a great deal of experience with over-calling protein identifications when stringent FDR thresholds are not maintained. The sensitivity to protein matching protocols can be illustrated with the results from the HUPO Human Plasma Proteome Project (HPPP). The original HPPP team paper¹⁴⁴ highlighted a “core dataset” of 3020 proteins with two or more peptide matches,

but clearly delineated a broad range of values with other criteria. In contrast, States et al.¹⁴⁵ published a uniquely stringent version of the same heterogeneous data utilizing Bonferroni-type adjustment for multiple comparisons with 889 protein identifications. In 2011, Farrah et al.⁴¹ published a Cedar scheme (**Figure S1** in the supporting information) for HPPP that demonstrated stepwise recognition of 1929 canonical proteins (1% protein-level FDR) + 236 possibly distinguished, totaling 2165 not subsumed; + 2507 subsumed = 4672 peptide-set unique; + 5686 indistinguishable = 9358 sequence-unique; + 10,102 identical = 19,460 exhaustive list (suitable for cross-checking a different canonical set to see whether the match was lost in the choice of a “representative protein”; see **Figure S1** in the supporting information). By 2014 the Human Plasma Proteome had grown to 4005 canonical proteins, as documented in the comparison of kidney, urine, and plasma proteomes¹⁰.

The Spanish chromosome 16 team developed a method using transcription data from public repositories (GEO^{146, 147}) obtained with cancer samples, cell lines, and healthy tissues to identify samples that showed enrichment for missing proteins [**poster 12**]. The data analysis showed that 2861 missing protein-coding genes were expressed at the mRNA level in at least one sample, and that the majority of the genes showed sample specificity. Their study confirmed that the missing proteins are typically shorter and of lower abundance than those that have been identified. Transmembrane, cytoskeleton, signal transduction, spermatogenesis, zinc finger domains, synapses, neurotransmitter activity, and olfactory transduction are enriched cellular functions among the missing proteins¹⁴⁸. All of these data will be available in the dasHPPboard webtool (<http://sphppdashboard.cnb.csic.es/>) [**poster 10**], which has the goal of creating a similar initiative for storing and accessing the processed data generated by the C-HPP projects in a manner similar to that of ENCODE^{46,47}.

To support the C-HPP initiative, Islam et al.¹⁴⁹ developed the Protannotator tool to provide extensive annotation of missing proteins. Protannotator consists of a generic pipeline incorporating bioinformatics and annotation tools to identify homologues and to map putative functional signatures,

gene ontology, and biochemical pathways. Sequential BLAST searches originally developed for chromosome 7¹⁵⁰ can be used to identify homologues from nonhuman mammalian proteins with strong protein evidence or homologues with validated human proteins. The Protannotator tool identified nonhuman mammalian homologues with protein evidence for 1271 missing proteins in other mammalian species, and 564 missing protein sequences were homologues to the reviewed human proteins. Functional annotations for the remaining missing proteins support the identification of possible biological sources and conditions under which the remaining missing proteins may be expressed. The tool also generates in silico proteotypic peptides, which facilitate the development of SRM assays. A search of these proteotypic peptides in ENCODE^{46, 47} revealed proteomic evidence for 107 missing proteins, with evidence for an additional 15 missing proteins using the data of a recent membrane proteomic study¹⁴⁹.

NeXtProt provides primarily web-based protein evidence information, but also enables retrieval of data in various output formats (HTML, JavaScript Object Notation [JSON] and XML) using the REST Application Programming Interface (www.nextprot.org/rest/). In addition, neXtProt provides “chromosome reports” on its ftp server to support C-HPP projects. At the workshop in Segovia, the neXtProt team announced the development of an advanced search engine based on SPARQL that will enable complex and powerful queries, including federated queries with external resources¹⁵¹.

Conclusions

The reduction of the proportion of missing proteins in the human proteome from 33% to 18% (or 15%) over the last four years shows the clear progress of the C-HPP, which is mainly due to the application of improved proteomics technology such as specific sample preparation (e.g., antibody-based enrichment and enrichment of hydrophobic peptides), the use of advanced spectrometers, the application of SRM and SWATH assays for missing proteins [**poster 3 and 14**], and the analysis of unusual human sample types [**posters 6, 7, 11, 13, and 15**]. As the results approach saturation of the

parts list for protein-coding genes, it will be ever more important to apply stringent FDR criteria to the claims of protein matches and to confirm the findings with orthogonal methods. “One-hit wonders,” especially of short peptides, and claims of matches in tissue or cell types without transcript expression or that have not previously shown evidence of such proteins with modern instruments should be viewed with skepticism. The quality of the spectra must be examined, keeping in mind that when Ezkurdia et al.¹³² examined the spectra for hundreds of olfactory receptor proteins claimed by Kim et al.⁷² and by Wilhelm et al.¹³¹, none survived scrutiny. Likewise, peptides with multiple matches may be more likely to represent known, highly expressed proteins with a single mutation or an RNA-edited site than a “missing protein.” The C-HPP has also encouraged analyses of amplicons (*cis*-regulated genes in specific chromosomal segments) and of protein families, as well as the recognition of proteins that are unlikely to be detected for the reasons outlined in **Figure 2**.

Proteogenomic analysis integrating data from genomics, transcriptomics, and proteomics is gaining momentum and results in an addition to the human proteome protein forms that arise from genetic variability, such as SAVs, RNA-editing, and alternative splicing. Proteogenomic technology now allows the routine study of these new protein forms in biological processes to unravel their roles in various diseases. Spectral libraries of synthetic peptides for almost all human proteins, together with the large number of antibodies generated by the HPA, permits the functional analysis of proteins and protein forms in biological experiments with complex designs. Bioinformatics support for the C-HPP has been largely developed during the last four years and has contributed to its success not only by reducing the number of missing proteins, but also in aiding the discovery of multiple new protein forms.

It is clear that work must still be undertaken to confirm the presence of the remaining missing proteins, which will become more and more challenging as the completion of the MS-based evidence of the human proteome on the gene basis is reached. C-HPP members are increasing their activities to find

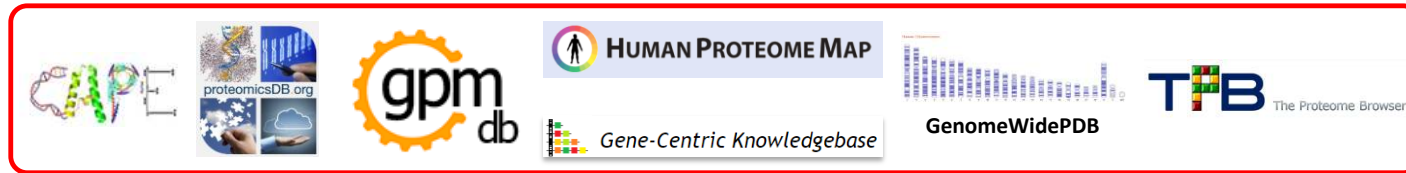
evidence for the remaining missing human proteins and to discover more and more complete sets of protein forms that reflect genetic variability and post-translational modifications.

The C-HPP posters presented at HUPO 2014 in Madrid (**Table 1**) are available online at the *Journal of Proteome Research* as supporting information, including the poster's abstract, and most of the oral presentations can be found at C-HPP Wiki (<http://c-hpp.webhosting.rug.nl/>).

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Unified Human Proteome Browser



Proteome Browsers

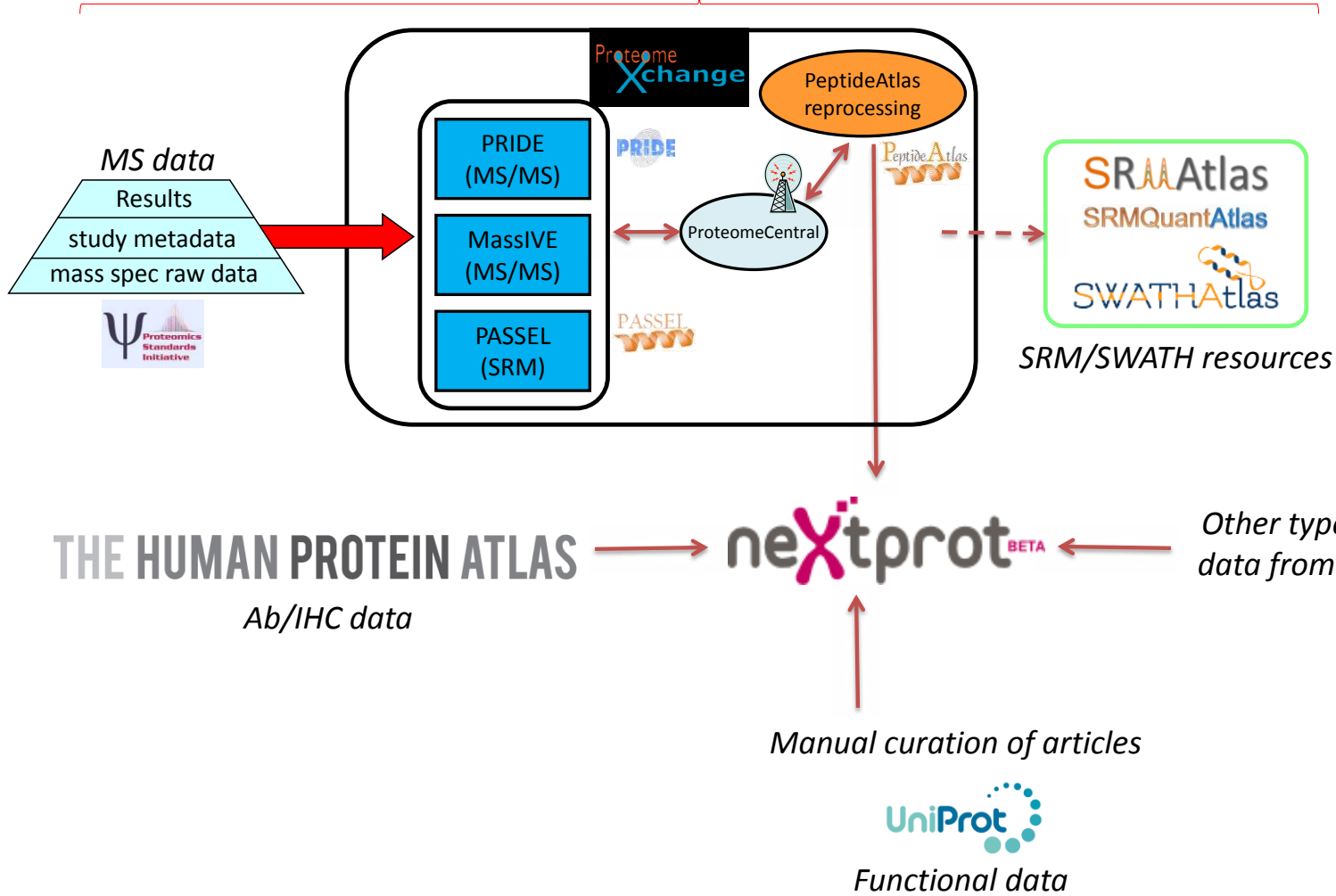


Figure 1. Bioinformatics resources to support the discovery, cataloging and browsing of protein part lists. Raw MS data acquired in different laboratories are deposited in the storage resources from the ProteomeXchange consortium. At present MS/MS datasets are fully supported by PRIDE and MassIVE, whereas SRM datasets are supported by PASSEL. Once is made publicly available, data in ProteomeXchange can be further used by many resources, for example reprocessing by PeptideAtlas using the Trans Proteomic Pipeline, or e.g. used by other Protein Browser resources. SRMAtlas, SRMQuanAtlas, SWATHAtlas with PeptideAtlas and other spectral libraries form the rich resources to develop or implement SRM assays. NeXtProt integrates data on proteins using 14 different resources and classifies proteins in 5 existence categories. C-HPP at HUPO 2014 in Madrid launched the Unified Human Proteome Browser Initiative to provide a unified view of the acquired proteogenomic data from a chromosome, biology and disease perspective.

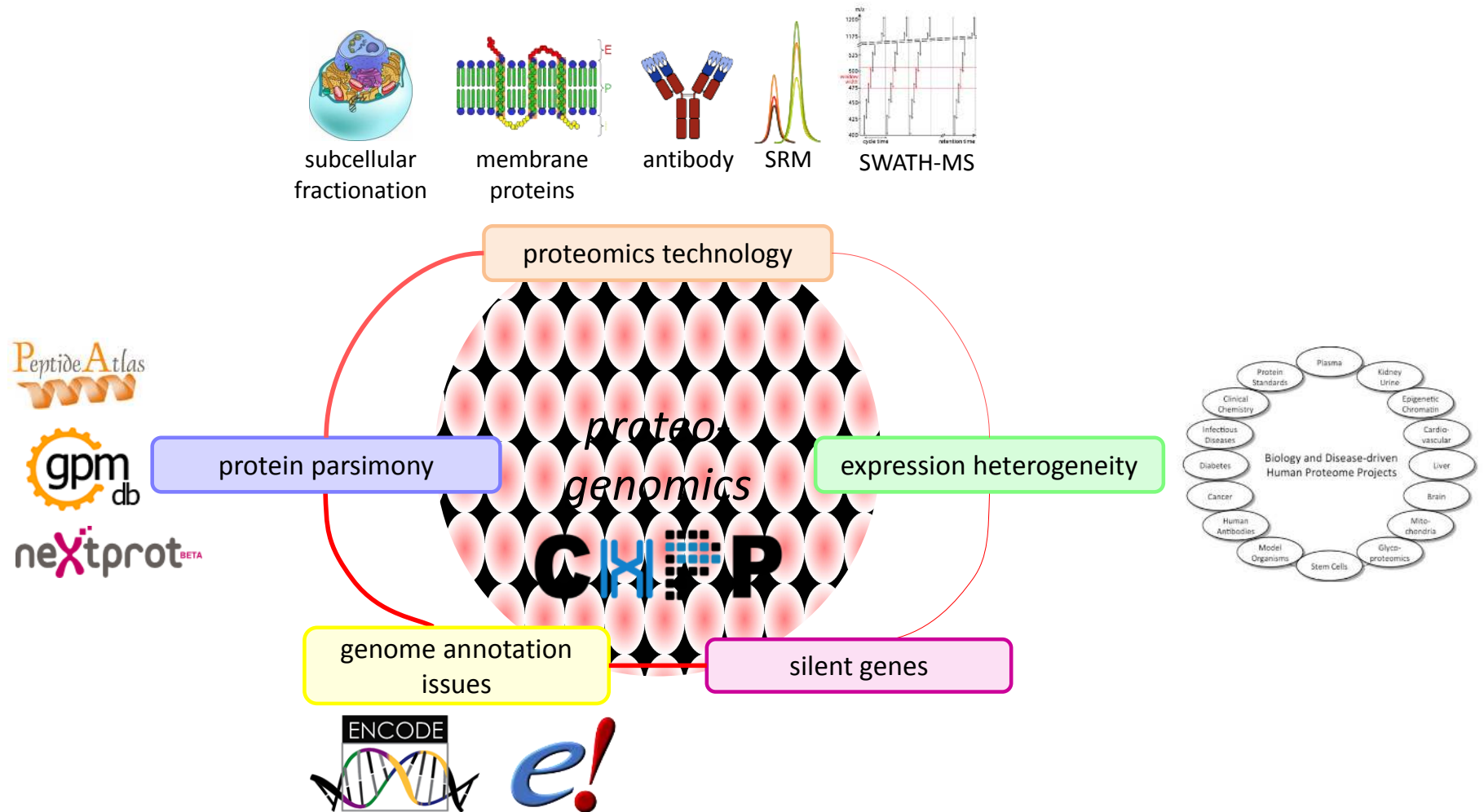


Figure 2. The five main reasons for proteins without evidence at the protein level (missing proteins) are: (1) current proteomics technology is not able to detect them due to uncovered chemical space of the applied mainstream analytical method, (2) expression heterogeneity of protein present only in rare and not yet analyzed samples, (3) silent genes present only in the genome, but never expressed, and (4) error in genome annotation, or (5) proteins missed or not counted due to parsimonious protein identification of shotgun proteomics database search

approach leading to simplification of protein representation of highly homologous proteins of the same protein family in databases such as PeptideAtlas, neXtProt, and gpmDB⁴⁰ or to large sequence variability such as immunoglobulins. Proteomics technology (1) can be improved with techniques such as subcellular fractionation or specific enrichment of membrane proteins, while low abundant proteins can be detected either with enrichment using monoclonal and polyclonal antibodies or/and using sensitive SRM and SWATH analysis. Expression heterogeneity (4) can be improved by joining forces with biology/disease driven research groups for example by enhancing collaboration between C-HPP and B/D HPP teams. Proteogenomic approach integrating genome, transcriptome with proteome data helps in general to (5) identify protein forms originating from genetic variability and (3) may correct for genome annotation errors. Use of multiple protease enhance protein coverage and can lead to distinction of highly homologous proteins (5) in main protein evidence databases. The most challenging group of missing proteins are silent genes (2) which are not normally expressed during the life cycle of an individual but can be activated by mutation, recombination, insertion elements, or other genetic mechanisms.



Figure 3. The number of chromosome 19 genes¹⁵² and the identified molecular entities at transcript and expression levels (mRNA and proteins) are illustrated as a proteogenomic analysis of *Glioma* stem cells addressed the challenges integrating genomics, transcriptomics and

proteomics data. Although, the figure presents the current status of chromosome 19, the number of “missing” consensus proteins and their alternative forms, including ASV, new ORFs and new SAVs, is proportionally similar of other human chromosomes.

Poster number	Title	Chr team	Main topic
1	Mining Missing Membrane Proteins from Lung Cancer Tissues and Cell lines	4	Proteomics technology: opening chemical space
2	CAPER 3.0: a scalable cloud-based pipeline for the data-intensive analysis of proteomic datasets	1, 8, 20	Bioinformatics: proteome browser
3	Chromosome 18: Master Proteome	18	Proteomics technology: lowering detection limit
4	GenomewidePDB v 2.0: Update on the transcriptomic and proteomic expression data with alternatively spliced products layered in a genome-wide manner	13	Bioinformatics: proteome browser
5	Development of improved subcellular fractionation procedures of the placental membrane proteins for discovering disease biomarkers and missing proteins	13	Proteomics technology: lower detection limit
6	Chromosome X	X	Missing protein: identification strategy
7	The Mitochondrial Human Proteome Project - MT-HPP	Mitochondria	Missing protein: identification strategy
8	Revisiting the Identification of Canonical Splice Isoforms through Integration of Functional Genomics and Proteomics Evidence from the Chromosome 17 Human Proteome Project	17	Exploring effect of genetic variability
9	Subcellular fractionation enhances Chromosome 16 Proteome Coverage	16	Proteomics technology: lowering detection limit
10	Proteogenomics Dashboard for the Human Proteome Project	16	Bioinformatics: proteogenomic data mining
11	Missing proteins in Chromosome 16 Spanish HPP	16	Missing protein: identification strategy
12	Transcriptomic profiling towards the localization of the missing proteins	16	Missing proteins: identification with proteogenomic
13	The Chromosome 19 Strategy to Characterize Novel Proteoforms and Missing Proteins Using ENCODE Resources	19	Missing protein: identification with proteogenomic
14	Targeting proteins of chromosome 16	16	Proteomics technology: lower detection limit
15	Dissecting Chromosome 16 proteome	16	Missing protein: identification strategy
16	Unified Human Proteome Browser Initiative	all	Bioinformatics: proteome

			browser
17	Expression of $\alpha v\beta 6$ integrin enhances both plasminogen and latent-transforming growth factor- $\beta 1$ dependent proliferation, invasion and ERK1/2 signaling in colorectal cancer cells	7	Proteomics technology: application
18	Overexpression of $\alpha v\beta 6$ integrin alters the colorectal cancer cell proteome in favor of elevated proliferation and a switching in cellular adhesion which increases invasion	7	Proteomics technology: application
19	Approaching the organellar brain proteome to understand the molecular basis of Schizophrenia	15	Protein quantification: application. Missing proteins.
20	Splice Variants in Aggressive Human Triple Negative Breast Cancer	17	Exploring effect of genetic variability
21	Revisiting the Identification of Canonical Splice Isoforms through Integration of Functional Genomics and Proteomics Evidence from the Chromosome 17 Human Proteome Project	17	Exploring effect of genetic variability

Table 1. List of posters presented at C-HPP poster session on 7 October 2014 at HUPO 2014 (Madrid), and used as second reference (poster number) in this paper.

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